IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicants: Thomas KOTHE, et al. Docket No.: COZ-0535

Serial No. 10/584,869 Examiner: Angela C. SCOTT

371 Date: April 20, 2007 Group Art Unit: 1796

Title: SETTABLE COMPOSITION CONTAINING LIME. A CEMENTITIOUS

COMPOSITION AND A POLYMER

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Vincent A. Cortese

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APRIL 8, 2010

(date)

APPELLANTS' BRIEF UNDER 37 C.F.R. § 41.37

To the Honorable Commissioner For Patents:

This is an appeal to the Board of Patent Appeals and Interferences (the "Board") from the final rejection set forth in the Office Action mailed December 9, 2009.

In accordance with 37 C.F.R. § 41.31, Appellants electronically filed the Notice of Appeal via EFS-Web on February 8, 2010.

The present appeal is of pending claims 1-20.

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1. Real Party in Interest

The owner of the present patent application is Construction Research & Technology GmbH, by virtue of an assignment from the Appellants. Construction Research & Technology GmbH is a limited liability company organized under the laws of the country of Germany, and is a subsidiary of BASF SE, a corporation organized under the laws of the European Union. The assignment for this application was recorded in the records of the Assignment Division of the United States Patent and Trademark Office (the "Office" or "USPTO") on August 14, 2006 at Reel/Frame 018098/0150.

2. Related Appeals and Interferences

In accordance with 37 C.F.R. § 41.37(c)(1)(ii), Appellants hereby inform the Board that there are no other prior pending appeals, interferences, or judicial proceedings known to Appellants, Appellants' legal representative, or Assignce which may be related to, directly affect or be directly affected by, or have a bearing on the Board's decision in the pending appeal.

3. Status of Claims

The present application was given a 35 U.S.C. § 371 date of April 20, 2007 after being initially submitted to the Office on June 29, 2006 with original claims 1-21. A Final Office Action for this application was mailed by the Office on November 24, 2008. A Request for Continued Examination for this application was filed with the Office on February 23, 2009. A Final Office Action for this application, subsequent to Appellants' Request for Continued Examination, was mailed by the Office on December 9, 2009.

Claims 1-20 are currently under final rejection and constitute the claims on appeal.

In accordance with 37 C.F.R. § 41.37(viii), appealed claims 1-20 appear in the Claim Appendix below.

4. Status of Amendments

A Final Office Action was mailed by the Office on December 9, 2009.

No amendments to pending claims 1-20 have been filed with the Office subsequent to the mailing date of the Final Office Action.

5. Summary of Claimed Subject Matter

Independent claim 1 is directed to a settable mixture comprising: (i) a water absorbing composition; and (ii) an aqueous emulsion of organic polymer; or (iii) dispersible organic polymer, wherein the water absorbing composition (i) contains inorganic ingredients which are capable to react with water and the water absorbing composition (i) contains at least 13 weight % lime and at least 5 weight % of a cementitious composition of which components form ettringite during the absorption of water and wherein the amount of the aqueous emulsion of organic polymer (ii) in relation to (i) is such as to provide a ratio of combined weight of polymer solids to combined weight of ingredients which are capable to react with water of from 0.5:1 to 10:1. and wherein the amount of dispersible organic polymer (iii) in relation to (i) is such as to give a ratio of combined weight of polymers to combined weight of ingredients which are capable to react with water of from 0.5:1 to 10:1. Specification at page 2, paragraph 4, through page 5, paragraph 1.

Dependent claim 5 is directed to a settable mixture as described above with respect to claim 1, wherein in respect to the formation of ettringite the water absorbing composition (i) contains a stoichiometric surplus of lime. Specification at page 5, paragraph 6, first sentence,

Dependent claim 13 is directed to a settable mixture as described above with respect to claim 1, wherein the water absorbing composition (i) contains at least 62% lime. Specification at page 5, paragraph 2.

Dependent claim 14 is directed to a settable mixture as described above with respect to claim 1, wherein the sum of the weight of the lime and the weight of the cementitious composition is 67 of 100 of the total weight of the water absorbing composition (i), and wherein the water absorbing composition (i) contains at least 62% lime. Specification at page 4, paragraph 5 (as amended) and page 5, paragraph 2,

Dependent claim 16 is directed to a settable mixture as described above with respect to claim 1, wherein the sum of the weight of the lime and the weight of the cementitious composition is 67 of 100 of the total weight of the water absorbing composition (i), and wherein in respect to the formation of ettringite the water absorbing composition (i) contains a stoichiometric surplus of lime. Specification at page 4, paragraph 5 (as amended) and page 5, paragraph 6, first sentence.

6. Grounds of Rejection to be Reviewed on Appeal

A. The 35 U.S.C. § 103(a) rejection of claims 1-4, 6-12, 15 and 17-20 as being unpatentable over U.S. Patent Application Pub. No. 2002/0161071 to Mills, et al., in view of U.S. Patent No. 6,514,334 B1 to Perry, et al. Appellants wish to note that, while the rejection on page 2 of the final Office Action mailed December 9, 2009 lists claims 1-4, 6-12, 15 and 17-20, the summary on page 1 of the final Office Action and the discussion of the rejection at pages 2-4 refers to all of claims 1-20. Appellants therefore assume for the purposes of this Brief that all of claims 1-20 have been rejected.

7. Argument

Claims 1-4, 6-12. 15 and 17-20 have been rejected under 35 U.S.C. § 103(a) as being unpatentable over U.S. Patent Application Pub. No. 2002/0161071 to Mills. et al. ("Mills"), in view of U.S. Patent No. 6,514.334 B1 to Perry, et al. ("Perry"). Appellants wish to note that, while the rejection on page 2 of the final Office Action mailed December 9, 2009 (the "Final Office Action") lists claims 1-4, 6-12, 15 and 17-20, the summary on page 1 of the Final Office Action and the discussion of the rejection at pages 2-4 refers to all of claims 1-20. Appellants therefore assume for the purposes of this Brief that all of claims 1-20 have been rejected.

While the Office admits that Mills does not teach a composition which contains at least 13 weight % lime, the Office has alleged that Perry teaches a cementitious composition where lime may be present in amounts greater than 62 weight %. Preliminarily. Appellants respectfully submit that the Office has failed to consider the arguments presented in Appellants' Response submitted August 8, 2008 and in Appellants' Response submitted February 23, 2009 regarding the difference between "lime" and the convention of referring to calcium oxide as a constituent oxide of a distinct composition. These arguments were previously presented with regard to the difference between lime and the "lime content" of calcium aluminate, but apply equally to the differences between lime and the "calcium oxide content" of a cementitious composition.

As described in H.F.W. Taylor, <u>Cement Chemistry</u>, <u>2d</u>, 1997, a copy of the pertinent portion of which is included as Appendix A, in the paragraph bridging pages 3 and 4, "[c]hemical formulae in cement chemistry are often expressed as sums of oxides; thus tricalcium silicate, Ca₃SiO₅, can be written as 3CaO·SiO₂. This does not imply that the constituent oxides have a separate existence within the structure." (emphasis added). This nomenclature is confirmed in Mills, at paragraph [0018].

Lime and calcium oxides which are constituent oxides of cementitious compositions are distinct compounds which react differently, and which must react differently in order to have their respective necessary effects on cementitious compositions, as required in the present application. The calcium oxide "content" of cementitious compositions is not available to react as lime, as it remains a part of the cementitious composition. If the lime "content" of a cementitious composition were to react as lime within the cementitious composition, there would

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be no calcium left to form the clinker phases necessary for the cementitious composition to act as a hydraulic binder. Further, it is well known in the art that the various oxides that form many cements, such as calcium oxide, alumina, ferric oxide and silica, are, by themselves, rather unreactive with respect to water, and thus a mere mixture of these oxides would not work as a hydraulic binder. The cement manufacturing process is necessary to transform these oxides into a crystalline form which is suitable for reaction with water and can therefore react as a hydraulic binder.

For these reasons, the 80-99% CaO content of composition (B) of Perry is not free to react as lime, as recited by the claims currently under appeal. It is therefore clear that a person of skill in the art would not look to Perry for a teaching of free lime content to modify Mills as alleged by the Office. Appellants therefore respectfully submit that the Office has failed to state a prima facie case of obviousness with regard to the combination of Mills and Perry, and request that the Board require that the rejection of claims 1-20 be withdrawn.

Further, Appellants respectfully submit that Mills and Perry are not properly combinable in order to establish a prima facie case of obviousness. MPEP § 2141.03(VI) states that "[a] prior art reference must be considered in its entirety, i.e., as a whole. including portions that would teach away from the claimed invention. W.L. Gore & Associates, Inc. v. Garlock, Inc., 721 F.2d 1540, 220 USPQ 303 (Fed. Cir. 1983), cert. denied, 469 U.S. 851 (1984)". MPEP at 2100-126 (emphasis in original). Further, MPEP § 2143.01(V) states that, "[i]f proposed modification would render the prior art invention being modified unsatisfactory for its intended purpose, then there is no suggestion or motivation to make the proposed modification. In reGordon, 733 F.2d 900, 221 USPQ 1125 (Fed. Cir. 1984)". MPEP at 2100-140.

The present specification discloses that, in order to achieve the desired high early strength provided by the presently claimed settable mixture, formation of Ettringite is necessary. At page 4, lines 22-26 (as amended), it is stated that "Ettringite is a calcium trisulphoaluminate having 32 molecules of water of crystallization and has the formula 3CaO.AL₂O₃.3CaSO₄.32H₂O. Ettringite is produced by the hydration of cementitious materials containing calcium aluminate

and <u>calcium sulphate</u>" (emphasis added). Mills contains a similar disclosure. Therefore, in order for the present settable mixture, as well as that of Mills, to be suitable for their intended purposes, the composition must contain calcium sulphate.

Perry does not disclose the use of calcium sulphate, and the composition of Perry cannot form Ettringite. Thus, if the composition of Perry, taken as a whole, were to be combined with Mills, the resulting composition would not contain calcium sulphate, as this component is absent in Perry, and therefore could not form Ettringite. The composition would not be able to obtain a high early strength. For these reasons, combining Perry with Mills renders Mills unsatisfactory for its intended purpose, and there is no suggestion or motivation to modify Mills according to Perry in order to increase the lime content of the Mills composition.

Furthermore, Mills discloses a maximum amount of 10% by weight of lime. See Abstract, paragraph [0010]. For this reason, there is no motivation in Mills to increase the amount of lime according to the teaching of Perry, as alleged by the Office.

Also for the above reasons, a person of skill in the art would not look to Perry for ways to modify Mills in order to obtain an even earlier high strength cementitious composition than that disclosed in Mills. Mills makes no disclosure of https://www.mills.com/bearly-strength, while the present specification shows that additional lime will provide a settable composition with an even earlier high strength than that of Mills. Perry provides no motivation to reduce the time needed for Ettringite formation by adding excess lime, because Perry does not contain the components necessary to form Ettringite. For all of these reasons, the Office has not established a prima facie case of obviousness.

Appellants submit that, since claim 1 is not taught or suggested by the combination of Mills and Perry, for the above reasons, claims 2-20, which depend from claim 1, are also not taught or suggested. See In re Fine, 837 F.2d 1071, 5 USPQ2d 1596 (Fed. Cir. 1988). ("If an independent claim is nonobvious under 35 U.S.C. 103, then any claim depending therefrom is nonobvious," MPEP § 2143.03 at page 2100-142.)

Specifically with regard to claims 5 and 16, the Office admits, at page 3 of the Final Office Action, that "Mills et al. does not teach that the water absorbing composition contains a

stoichiometric surplus of lime, i.e., an amount of lime that is not included in the cttringite forming reaction." The Office alleges, however, that "Perry et al. teaches adding up to 30 weight percent hydrated lime to the cementitious mixture comprising compositions A and B. Adding this lime after the composition has been formed would preclude it from participating in the reaction." Appellants respectfully submit that this assertion is technically incorrect.

Claims 5 and 16 of the present application are directed to compositions of claim 1 or claim 2, "wherein in respect to the formation of ettringite the water absorbing composition (i) contains a stoichiometric surplus of lime."

Perry discloses a process for making a cementitious mixture which includes mixing granular components A and B (Abstract). The Office alleges that the reaction of components A and B will be complete once components A and B are mixed. This is not true. Components A and B, when mixed, form a "homogenous cementitious mixture" (col. 6. lines 40-41). However, as is well known in the art of cementitious compositions, components A and B will not react until water is added to the homogenous mixture. Therefore, the addition of up to 30 weight percent lime (col. 6, lines 44-45) does not teach a stoichiometric surplus in any sense, because the reaction has not yet begun and the required amount for reaction not indicated, when the lime is added to the homogenous mixture of Perry.

For these reasons, Perry gives no indication that any additional lime added to the cementitious mixture of Perry is more than what will be required or consumed in the cementitious reaction that will occur once water is added to the cementitious mixture of Perry. Further, Perry makes no mention of providing lime at a stoichiometric surplus, or that there is any desire to increase the speed of the reaction of components A and B once water is added to the cementitious mixture. Therefore, there is no teaching, suggestion, or motivation in Perry to add a stoichiometric surplus of lime to the cementitious mixtures of Mills and/or Perry.

For this reason, Appellants respectfully submit that the combination of Mills and Perry fails to teach or suggest the subject matter of claims 5 and/or 16 of the present application.

Specifically with regard to claims 13 and 14, the Office alleges, at page 2 of the Final Office Action with regard to Perry, that "assuming that composition B is made of 90% lime, lime would be in the composition in an amount greater than 62 weight %".

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Claims 13 and 14 of the present application are directed to compositions of claim 1 or claim 2, "wherein the water absorbing composition (i) contains at least 62 weight % lime."

As discussed above, the alleged "lime" of composition B, being part of a complex oxide structure and not having a separate existence within the structure, is not free to react as lime reacts in the present application. Perry discloses a maximum amount of 30 weight percent lime (col. 6, line 45). Therefore, the combination of Mills and Perry fails to teach or suggest the subject matter of claims 13 and/or 14 of the present application.

For all of the above reasons, Appellants respectfully request that the Board require that the rejection of claims 1-20 be withdrawn.

Conclusion

Appellants have addressed the instant rejections presented in the Final Office Action with respect to independent claim 1 and dependent claims 5, 13, 14 and 16 in particular, and have distinguished the applied references as discussed above. It is therefore deemed unnecessary to address the Office's specific allegations regarding the remaining dependent claims. Appellants therefore traverse these allegations, and do not concur with the same either explicitly or implicitly by not refuting each individually.

Appellants submit that the remarks presented hereinabove overcome all the existing 35 U.S.C. § 103(a) rejections of all pending claims. Appellants respectfully request the Board to reverse the final rejections of these claims, and to require the Office to issue a formal notice of allowability of claims 1-20 over the art of record.

Respectfully submitted.

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Attorneys for Appellants Date: Upril 8, 2010

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8. Claims Appendix

In accordance with 37 C.F.R. § 41.37 (c)(1)(ix), the claims on appeal are as follows:

- 1. (Previously Presented) A settable mixture comprising
 - (i) a water absorbing composition and
 - (ii) an aqueous emulsion of organic polymer or
 - (iii) dispersible organic polymer,

wherein the water absorbing composition (i) contains inorganic ingredients which are capable to react with water and the water absorbing composition (i) contains at least 13 weight % lime and at least 5 weight % of a cementitious composition of which components form ettringite during the absorption of water and

wherein the amount of the aqueous emulsion of organic polymer (ii) in relation to (i) is such as to provide a ratio of combined weight of polymer solids to combined weight of ingredients which are capable to react with water of from 0.5:1 to 10:1, and

wherein the amount of dispersible organic polymer (iii) in relation to (i) is such as to give a ratio of combined weight of polymers to combined weight of ingredients which are capable to react with water of from 0.5:1 to 10:1.

- (Previously Presented) A settable mixture according to claim 1, wherein the sum of the
 weight of the lime and the weight of the cementitious composition is 67 to 100 of the
 total weight of the water absorbing composition (i).
- (Previously Presented) A settable mixture according to claim 1, wherein the water absorbing composition (i) contains at least 25 weight % lime.
- (Previously Presented) A settable mixture according to claim 1, wherein the cementitious composition contains calcium aluminate and calcium sulfate.
- (Previously Presented) A settable mixture according to claim 1, wherein in respect to the formation of ettringite the water absorbing composition (i) contains a stoichiometric surplus of lime.

6. (Previously Presented) A method of applying a coating to a surface which method comprises forming a settable mixture according to claim 1, containing a water absorbing composition (i) and an aqueous emulsion of organic polymer (ii) and putting the settable

mixture on the surface to form a coating.

7. (Previously Presented) A method of applying a coating to a surface which method comprises forming a settable mixture according to claim 1, containing water absorbing composition (i) and dispersible organic polymer (iii), combining the settable mixture with water and putting the mixture on the surface to form a coating.

 (Previously Presented) The coating made according to the method which is defined in claim 6.

9. (Previously Presented) A method of applying a coating to a surface which method comprises forming a settable mixture according to claim 1, containing a water absorbing composition (i) and an aqueous emulsion of organic polymer (ii) and putting the settable mixture on the surface to form a coating, wherein the coating comprises a rock support means.

10. (Previously Presented) A method of applying a coating to a surface which method comprises forming a settable mixture according to claim 1, containing a water absorbing composition (i) and an aqueous emulsion of organic polymer (ii) and putting the settable mixture on the surface to form a coating, wherein the coating comprises a waterproofing means.

11. (Previously Presented) The settable mixture according to claim 1 wherein the amount of the aqueous emulsion of organic polymer (ii) in relation to (i) is such as to provide a ratio of combined weight of polymer solids to combined weight of ingredients which are capable to react with water of from 1: 1 to 4: 1.

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12 (Previously Presented) The settable mixture according to claim 1 wherein the amount of dispersible organic polymer (iii) in relation to (i) is such as to give a ratio of combined weight of polymers to combined weight of ingredients which are capable to react with

water of from 1: 1 to 4: 1.

13 (Previously Presented) The settable mixture according to claim 1 wherein the water

absorbing composition (i) contains at least 62 weight % lime.

14. (Previously Presented) The settable mixture according to claim 2 wherein the water

absorbing composition (i) contains at least 62 weight % lime.

15. (Previously Presented) The settable mixture according to claim 2 wherein the

cementitious composition contains calcium aluminate and calcium sulfate.

16. (Previously Presented) The settable mixture according to claim 2 wherein in respect to

the formation of ettringite the water absorbing composition (i) contains a stoichiometric

surplus of lime.

17 (Previously Presented) The method of applying a coating to a surface according to claim

6 wherein the coating is at least 2 mm in thickness.

18. (Previously Presented) The method of applying a coating to a surface according to claim

7 wherein the coating is at least 2 mm in thickness.

19. (Previously Presented) The coating made according to the method which is defined in

claim 7.

(Previously Presented) A method of applying a coating to a surface which method

comprises forming a settable mixture according to claim 1, containing water absorbing composition (i) and dispersible organic polymer (iii), combining the settable mixture with

water and putting the mixture on the surface to form a coating, wherein the coating

comprises a rock support means or a waterproofing means.

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9. Evidence Appendix

Appendix A: Taylor, H.F.W., <u>Cement Chemistry</u>. 2nd Edition, Thomas Telford Publishing, 1997. pages 1-5.

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APPENDIX A

Cement chemistry

2nd edition

Emeritus Professor of Inorganic Chemistry University of Aberdeen H.F.W. Taylor

Thomas Telford

> Judished by Thomas Telford Publishing, Thomas Telford Services Ltd., Heron Quay, London E1441D First published 1990 by Academic Press Thomas Telford edition published 1997

Preface

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The previous edition of this book, published by Academic Press in 1990, is both our of print and our of clear, and when Thomas Telford invited me to prepare a new edition I was edigited to comply. I am most gardell obth to the man of the copyright made both to there and to Academic Press, whose release of the copyright made both to the real of the Academic Press, whose release of the copyright made the present edition possible.

With some dash, which extensing of the principal idies not duminate exactan used in bailing and off or experiently, it is directed parally to these whose transparent in including states extense or related disciplane, Emphasis is placed throughout on the underlying science make than on proteinal papelasisms, which are well covered in online works. The exempts considered full ston let cangery of hyboratic and funded with some transparent and the parallel can central Canador in the destination of the transparent is britten exerted. Canador is the destination which the result important is britten exerted. Canador is the destination which the result important is britten exerted. Canador is destination which the result important is britten of careful. Canador is destination which the contributions of britten canador in the processor has been accorded in proposite in british with which the processor has been conveniently of britten corner where which the processor has been conveniently of britten for a way to any which the processor has been conveniently of the trained with and of special uses for cements. Chapter 12 deals with chemical and microstructural aspects of concrete, including ones that affect its durability or with the chemistry of other types of cement, of admixtures for concrete

1000 new contributions annually. The output of the seven years since the previous edition of this book appeared is reflected in the interessed number of references to the literature, which is approximately 1300 compared with 1300 in that edition. Of necessity, coverage in the present book has been selective, but it is hoped that the most important contributions up to mid-1996 have been covered. The advances in some parts of the subject have been greater than in others, and this is reflected in the differences between this and the previous edition; some sections have been totally rewritten whereas others have changed relatively little.

As one who has seen the subject develop over a period of nearly 50 The literature of cement chemistry is voluminous; the abstracting journal, Cements Research Progress, has for some years listed around rears, I am highly aware of the problems that those entering the subject imit its service life.

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Typesei by The Alden Group, Oxford Printed in Great Britain by Redwood Books, Trowbridge, Willakire

Portland cement and its major constituent phases

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Appendix: Calculated X-ray powder diffraction patterns for tricoleium silicate and clinker phases

References

Coment chemistry

.1 Introduction

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about the same under comparable conditions.

Aluminate constituce 5-10% of most anomal Portland cement clinters. It is tricalcium aluminate (CayAl-Q), substantially modified in composition and sometimes also in structure by ionic aubstitutions.

75 ×

Sament chemistry

tion by variation in AI/Fe ratio and ionic substitutions. The rate at which it reacts with water appears to be somewhat variable, perhaps due to differences in composition or other characteristics, but in general is high initially and low or very low at later ages. it reacts rapidly with water, and can cause undestrably rapid setting unkets a set-controlling agent, usually gypsum, is added. Ferrite makes up 5–15% of normal Pretland cement clinkers. It is tetracalcium aluminoferrite (Ca.AIFeOs), substantially modified in composi-

Types of Portland cement

The great majority of Portland cements made throughout the world are designed for general constructional use. The standard specifications with which such cements must comply are similar, but not identical, in (42.1) the internal T-84-year late of (40.2) the minum. T-84-year late of Popul and IP Portland centers in the ASTA (American Society for Traiting, and Martically September 10-10. Traiting, and Martically September 10-10. Traiting, and Corners (OPC) in former Birlish, standards. Throughout this book, the term orderary Portland centers it and to distinguish web general-purpose emensis (opo other types of Portland corners, which are made in maller.) all countries and various names are used to define the material, such as Class 42.5 Portland cement in current European and British standards

quantics for expectal present, its general, based partly on chemical Standard specifications are, in general, based partly on chemical emposition of publical properties; such as specific autice area, and partly on performance tests, such as setting time or comprastive acrossip processing the standard conditions. The control of MgO's is usually finded to 4-2%, bestage quantics of this component in stances of about The first own central springles in the control of t Other upper limits of composition widely used in specifications relate to matter fasoluble in dilute ueid, and loss on ignition. Many other minor components are limited in content by their effects on the manufacturing process, or the properties, or both, and in some cases the limits are defined in specifications.

Rapid-hardening Portland cements have been produced in various ways, such as varying the composition to increase the alite content, finer grading of the clinker, and improvements in the manufacturing process,

Concludes on stills because the mean of formulae of compounds can be used to denote offerent fairness on mean special this spaties are species by Co.C. and Mp. Oher and elsebate, obtained a mean special to desir tig, action words, important ands, then, performs all generally be used for places, and formulae of the CO.M. Mp. Compounds. Mirral interest or greated formulae (e.g. or other) just one used for compounds.

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Portland cament and its major constituent phases

the latter at practic in consequency proving in its recent in this in-manch on contract by militar delining. The reaction involves the Ad-D constituing, plants in the inherited current, and in sultimerships prelated carentis its effects are related by detersing the proportion of the stummines plants, arounds not some time its eithered by descrizing the ratio of Ad-D, to the two meantable, in either the description around great present and the pure venture in Visit and relating prefated means are stell Type V transmitted. May sultime relating formula demonst a real tall Type V transmitted of Ad-D, to Pd-D, and than represent representations to sultime resisting formula or means the propriet entering composition of partial or prefated and than represent representations are composition to sultime resisting formula or means the toronty and recomposition or sultime resisting formula or means. The toronty aid to color of Perdand carent Destructive expansion from reaction with sulfates can occur not only if

free from FrQ-5 and other components; such as Mr₂O₅, that combine to the colour. The effects of these components are therefore usually minimized by producing the clinical under slightly reducing conditions with any prapel questions. The clinical conditions are such as the conditions glass may be formed. is due to the ferrite, formation of which in a white cement must thus be avoided. It is impracticable to employ raw materials that are completely

moderate sulfate attack, and is widely used in general construction work. Heat evolution can also be decreased by parfully replacing the centent by Hyazh (pulverized nel ast, pill or other materials (Chapter) y, and this is today a widely used audition. The specialized requirements of oil well centions are discussed in Section 11.8. The reaction of Portland cement with water is exothermic, and while this can be an advantage under some conditions because it accelerates hardening, it is a disadvantage under others, such as in the construction of large dams or to the lining of oil wells, when a cement stury has to be pumped over a large distance under pressure and sometimes at a high emperature. Slower heat evolution can be achieved by coarser grinding, and decreased total heat evolution by lowering the contents of alite and aluminate. The ASTM specifications include definitions of a Type 11 or moderate heat of hardening coment, and a more extreme Type IV or 'low heat' coment. The Type II coment is also suitable for conditions exposed to

Chemical formulae in coment chemistry are often expressed as sums of oxides; thus trienterum silicate. CaySiO₂, can be written as 3CaO-SiO₂. This does not imply that the coveriment does not imply that the constituent oxides have any separate Cament chamical nomenclature and other abbraviations

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Cement chemistry

commonte order to the formation of the formation of the Commonte order to single elercy, such as C for C 50 × 50 × 50 C C A, S(O, thus becoming C,S. This system is often combined with orthodox chemical notation whith an elemental equation, e.g. (1,1) 30.00+50.0.—C.S. (1,1)

 $3C_2O + S(O_2 - C_3S)$ (1.1) or even within a single formula, as in $C_1A_7 - CaE_2$ for $Ca_1_2A|_{A_2}O_2F_2$. The abbreviations most widely used are as follows.

 $T=\overline{110}$, $P=P_2O_3$ $H=H_2O$ $\tilde{C}=CO_3$ The formulae of the simple oxide phases (e.g. CaO) are usually written in bill. Other abbreviations and units used in this book are as follows. H. The Photogram, A. H. The Photogram, A. H. The Photogram, BE a backstratered electron. BE a backstratered electron for the A. offercold internal tradyies. Best a descript of the factors profess price as 1945. ISCA estimate them to perturbe to the contraction of the profession of

113.2 Materials
C-S-H = poorly crystalice or amorphous calcium sileate hydrate of
the profit composition. Ggbs = promit granulated hasfurmeet stage.
Hsp = hardened senten passe, Fit = pulverized tied sah (pissk).

11.132 Properties or reactions
AR external ratio (chimae modulu), ASR = atkali silica reaction.
AR external ratio (chimae modulu), ASR = atkali silica reaction.
SN = silica ratio (chimae modulu), C. = analyte (chimae ratio (chimae)), C. = analyte (chimae ratio (chimae)), C. = analyte (chimae ratio), C. of chimae (chimae), C. = analyte (chimae), C. of chimae (chimae), C. analyte (chima

Nexy—rece x, v). r+, r v, v = 1, r v, optic sign and optic extent angle.
11.3.4 Pressure units
11.3.4 Pressure units
11.3.4 Pressure units
10.198 kg cm = 1, 87 atm = 7300 torr = 145.0 bin = 2 = 10.198 kg cm = 1.

11.2 Allte
11.2.1 Polymorphism and crystal structure
On bring heatel, pure C₃S undergoes a series of reversible phase
Intantions, which have been detected by a combination of DTA,

Portland coment and its major constituent phases high-temperature Ight microscopy (B1,G1, M3-M5,R1,R2,Y1);

promission distant, set on the interpoparion of pathitisms (not, the form persent si come manyanta numbic appelations on N to or M or as a form of the control of the contr

of disorder. The structured inferences between the job/mosphy affect the considiration of the Ce² box and this Contain of the Strip terrelated, to see the different conclusions of the conclusion contributed and the speed site that coordination contributed and the speed site that coordination contributed with the conclusion contributed and the speed site that coordination contributed and the speed site that coordination contributed and the speed site of the coordination contributed and the speed site of the coordination contributed and supports the speed site of the site could be repeated at Toolshead the Ordination of the Ca norm in one of the site could be repeated at Toolshead the other is to get Stefan are commute, and conducted and the structured are structured as the structure of the Ca; it is AGS in the R popuration by Stefan in the Ordination of the copyen stoms in possibly more constrained and the speed structured and the structured on the structured in the structured on the structured in the structured on the

incenter with those of culcions.
Table 11 gives the crystal data for the C5 polymorphis that have been oblighted single single-styral methods. The fine-tune constitute stellars are made and the crystal stellars are not a Table 11 diene be unit cit of a monocibied many be equivalent to ench of Table 11 diene be unit cit of a monocibied many between the control of Table 11 diene be unit cit of a monocibied many that the control of the cit of the control of the cit of the control of the cit of t

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10. Related Proceedings Appendix

None.